(19) World Intellectual Property Organization

International Bureau



(43) International Publication Date 24 June 2004 (24.06.2004)

PCT

(10) International Publication Number WO 2004/053895 A1

(51) International Patent Classification7:

H01B 3/44

(21) International Application Number:

PCT/EP2003/011905

(22) International Filing Date: 27 October 2003 (27.10.2003)

(25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

02027860.2

12 December 2002 (12.12.2002)

(71) Applicant (for all designated States except US): BORE-ALIS TECHNOLOGY OY [FI/FI]; P.O. Box 330, FIN-06101 Porvoo (FT).

(72) Inventors; and

(75) Inventors/Applicants (for US only): FAGRELL, Ola [SE/SE]; Gullkragevägen 5, S-444 45 Stenungsund (SE). NILSSON, Ulf [SE/SE]; Koppersvägen 42, S-444 42 Stenungsund (SE).

Agent: KADOR & PARTNER; Corneliusstrasse 15, 80469 Munich (DE).

(81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

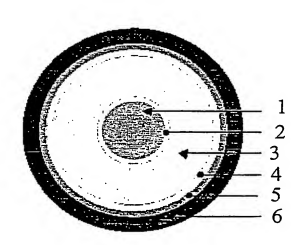
(84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

with international search report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: COAXIAL CABLE COMPRISING DIELECTRIC MATERIAL



(57) Abstract: The present invention relates to a coaxial cable comprising a dielectric layer which comprises as a component (A) a propylene homo- or copolymer having strain hardening behaviour and to the use of propylene homo- or copolymer having strain hardening behaviour for the production of a dielectric layer of a coaxial or triaxial cable.

WO 2004/053895 A1



Coaxial Cable Comprising Dielectric Material

The present invention relates to a coaxial or triaxial cable, in particular to a coaxial high radio frequency cable, comprising a dielectric layer, and to a dielectric material for use in a coaxial or triaxial cable.

A coaxial cable is defined to comprise one centre conductor and one outer concentric conductor and a triaxial cable is defined to comprise one centre conductor and two outer concentric conductors with an isolating layer separating them. Usually, these cables are protected with an outermost jacket.

In the following, where reference is made to coaxial cables also triaxial cables should be included.

In a coaxial cable the diameter of the dielectric material is typically above 1 mm. In radio frequency cables the diameter of the dielectric usually varies between 4 mm and 52 mm.

For the transfer of radio frequency signals e.g. in antenna systems of base stations of mobile phone networks, the use of coaxial cables is common in the art.

Typically, radio frequency coaxial cables are used as feeder or radiating cables. Feeder cables are used in the high power transmission from the power amplifier stage of a radio transmitter to the radiating antenna element or connection of a receiving antenna to the input stage of a radio receiver, or a combination of similar signal paths. An example of such an application is found at the base stations of mobile phone networks. Another application is in the radio shadow areas of said mobile phone systems such as tunnels, cellars, etc., where this type of cable can be used as the radiating element when provided with a perforated leaky outer conductor. The coaxial cables are useful also in community antenna television (CATV) systems in which the transmitted signal conveys both analogue and digital television pictures, as

well as on the subscriber lines of modern telephone systems (access networks) which use coaxial cables as the transmission medium in the transfer of wideband information.

A typical coaxial cable comprises an inner conductor made of copper or aluminium, a dielectric insulation layer made of a polymeric material, and an outer conductors made of copper or aluminium (see Fig. 1). Examples of outer conductors are metallic screens, foils or braids. Furthermore, in particular when polyethylene is used for the dielectric layer, the coaxial cable comprises a skin layer between the inner conductor and the dielectric layer to improve adherence between inner conductor and dielectric layer and thus improve mechanical integrity of the cable.

An important requirement for the dielectric layer of coaxial cables is that the attenuation of the signal should be as small as possible. Therefore, today said polymeric dielectric layer, typically polyethylene, is usually expanded by chemical or physical foaming to a level of up to 75 vol% or more.

However, due to the high degree of expansion typically used it is required for high frequency RF communications that the polymeric material used for the dielectric layer shows superior mechanical properties for the melt upon expansion to obtain closed and even cell structure.

For example, from US 6,130,385 it is known to use a blend of a low density polyethylene (LDPE) and medium density polyethylene (MDPE) for expandable dielectric layers of coaxial cables which shows good mechanical properties upon expansion. However, as today radio frequency cables tend to be used at ever increasing frequencies of up to several GHz it is a drawback of this dielectric layer material that the attenuation of the signals caused by the dielectric layer worsens with increasing frequency.

Furthermore, these cables have the disadvantage that the dielectric layer has to increase in thickness if the cable is used at higher frequencies and high power of the signal as required by the mobile phone networks of today and in future.

Accordingly, it is an object of the present invention to overcome the draw-backs of the above described techniques and to provide a coaxial or triaxial cable, especially for the transmission of high radio frequency signals, comprising a dielectric layer which is having a low attenuation, especially at higher radio frequencies and, at the same time, good mechanical properties of the melt so that a high degree of foaming can be achieved.

It has now surprisingly been found that such a coaxial or triaxial cable can be obtained if it comprises a dielectric layer which comprises polypropylene which has been modified in a particular way.

Accordingly, the present invention provides a coaxial and a triaxial cable comprising a dielectric layer which comprises as a component (A) a propylene homo- or copolymer having a strain hardening behaviour.

With the inventive cable the above-mentioned objects of the invention are achieved. In particular, the cable is showing an improved attenuation of the signal, especially at higher radio frequencies. It is believed that the improvement in attenuation is due to the particular behaviour of the so-called loss- or dissipation factor ($\tan \delta$) of the propylene homo- or copolymer used in the dielectric layer. This loss-factor has been found to be the most important influence factor for the attenuation behaviour of the dielectric layer.

The improved electrical properties of the inventive material enable higher operating frequencies and/or reduction in total cable thickness.

Due to the improved mechanical properties of the melt of the dielectric layer it is possible to obtain a high degree of expansion which also contributes to the good attenuation properties of the inventive cable.

It is a further advantage of the inventive cable that due to the improved mechanical properties of the dielectric layer a skin layer between the inner conductor and the dielectric layer can be omitted.

As polypropylene can withstand higher temperatures than polyethylene, the cable can be operated at a higher conductor temperature and therefore allows

WO 2004/053895 PCT/EP2003/011905

the transmission of signals with higher power rating and/or at higher frequencies.

The inventive cable can advantageously be used in all applications requiring the transfer of a radio frequency signal, especially at higher frequencies, whether digital or analogue. In particular, the cable can be used as feeder or radiating cable in mobile phone networks.

Propylene homo- and copolymers having strain hardening behaviour can be produced by a number of processes, e.g. by treatment of the unmodified propylene polymer with thermally decomposing radical-forming agents and/or by treatment with ionising radiation, where both treatments may optionally be accompanied or followed by a treatment with bi- or multifunctionally unsaturated monomers, e.g. butadiene, isoprene, dimethylbutadiene or divinylbenzene.

Further processes may be suitable for the production of the modified propylene polymer, provided that the resulting modified propylene polymer meets the characteristics of strain hardening behaviour, which is defined in the Examples Section below.

Examples of said modified propylene polymers showing strain hardening behaviour are, in particular:

- polypropylenes modified by the reaction of polypropylenes with bismaleinmido compounds in the melt as e.g. described in EP 0 574 801 and EP 0 574 804,
- polypropylenes modified by the treatment of polypropylenes with ionising radiation in the solid phase as e.g. described in EP 0 190 889 and EP 0 634 454,
- polypropylenes modified by the treatment of polypropylenes with peroxides in the solid phase, see e.g. EP 0 384 431, or in the melt, see e.g. EP 0 142 724,

- polypropylenes modified by the treatment of polypropylenes with multifunctional, ethylenically unsaturated monomers under the action of ionising radiation as described e.g. in EP 0 678 527,
- polypropylenes modified by the treatment of polypropylenes with multifunctional, ethylenically unsaturated monomers in the presence of peroxides in the melt as described e.g. in EP 0 688 817 and EP 0 450 342.

The modified propylene polymers having strain hardening behaviour are preferably prepared by

- a) mixing a particulate unmodified propylene polymer, which comprises
 - al) propylene homopolymers, preferably propylene homopolymers with a weight average molecular weight Mw of 500,000 to 1,500,000 g/mol, and/or
 - a2) copolymers of propylene and ethylene and/or alpha-olefins with 4 to 18 carbon atoms, or of mixtures of such copolymers,

with from 0.05 to 3 wt%, based on the polyolefin composition used, of acyl peroxides, alkyl peroxides, hydroperoxides, peresters and/or peroxycarbonates as free-radical generators capable of thermal decomposition, if desired diluted with inert solvents, with heating to 30 - 100°C, preferably to 60 - 90°C,

- b) sorption of bifunctional unsaturated monomers by the particulate propylene polymer at a temperature T (°C) of from 20 to 120 °C, preferably of from 60 to 100 °C, where the amount of the absorbed bifunctional unsaturated monomers is from 0.01 to 10 wt%, preferably from 0.05 to 2 wt%, based on the propylene used, and then
- c) heating and melting the particulate polyolefin composition in an atmosphere comprising inert gas and/or the volatile bifunctional monomers, from sorption temperature to 210°C, whereupon the free-radical generators capable of thermal decomposition are decomposed and then

- d) heating the melt of to 280°C in order to remove unreacted monomers and decomposition products,
- e) agglomerating the melt in a manner known per se.

Usual amounts of auxiliary substances, which may range from 0.01 to 1.5 wt% of stabilizers, 0.01 to 1 wt% of processing aids, 0.1 to 1 wt% of antistatic agents, 0.2 to 3 wt% of pigments and up to 3 wt% of alpha-nucleating agents, in each case based on the sum of the propylene polymers, may be added before step a) and/or e) of the method and/or before or during step c) and/or d) of the above described method.

The particulate unmodified propylene polymer may have the shape of powders, granules or grit with grain sizes ranging from 0.001 mm up to 7 mm.

The process for producing the modified propylene polymer preferably is a continuous method, performed in continuous reactors, mixers kneaders and extruders. Batchwise production of the modified propylene polymer, however is feasible as well.

Preferably volatile bifunctional monomers are absorbed by the particulate propylene polymer from the gas phase.

Practical sorption times τ of the volatile bifunctional monomers range from 10 to 1000 s, where sorption times τ of 60 to 600 s are preferred.

The bifunctional unsaturated monomers, which are used in the process for producing the modified propylene polymers preferably are C_4 - to C_{10} -dienes and/or C_7 - to C_{10} -divinyl compounds. Especially preferred are butadiene, isoprene, dimethyl-butadiene or divinylbenzene.

Preferably, the propylene homo- or copolymer having strain hardening behaviour has a melt flow rate of 0.1 to 25 g/10min at 230°C/2.16kg.

In a preferred embodiment of the present invention, the dielectric layer of the coaxial cable further comprises as a component (B) a medium or high density ethylene homo- or copolymer and/or a non-strain hardening behaviour propylene homo- or copolymer.

Medium density polyethylene typically has a density of 926 to 940 kg/m³ according to ASTM D 1248, and high density polyethylene typically has a density of 940 to 960 kg/m³.

If component (B) comprises polyethylene, it is preferred that it said polyethylene has medium density.

It is, however, preferred that component (B) comprises a non-strain hardening behaviour propylene homo- or copolymer, i.e. a polypropylene which after its production has not been modified to show strain hardening behaviour.

With the incorporation of said component (B) into the dielectric layer the mechanical properties and, in particular, the attenuation behaviour of said layer is further improved.

Further preferred, component (B) of the dielectric layer of the inventive coaxial cable comprises a clean-polypropylene.

Clean-polypropylene as used herein is defined to be a propylene homo- or copolymer, preferably a propylene homopolymer or ethylene copolymer having a catalyst residue less than 50 ppm, preferably less than 5 ppm, measured by ICP, an ash content below 100 ppm, preferably below 30 ppm, and a chloride content less than 5 ppm, preferably less than 1 ppm.

The catalyst residue is measured by determining of the amount of one or more elements present in the catalyst, usually Al, in a polypropylene sample by means of ICP, for example using a Plasma 40 Emission Spectrometer from Perkin-Elmer. Before the measurement, the polymer sample is brought into a soluble form, e.g. by careful burning of the sample at about 600°C, addition of Li₂CO₃ and NaJ, further heating to about 1000°C and dissolving the cooled sample in nitric acid solution.

The ash content is determined by ashing a polypropylene sample at 1000°C e.g. in a muffle furnace and weighing the rest.

The chloride content of a polypropylene sample is determined on the basis of X-ray fluorescence (XRF) spectrometry, e.g. by using an X-ray fluorescention Philips PW 2400.

Preferably, the clean-polypropylene is produced in a slurry process.

An example of clean-polypropylene as mentioned above is described, for example, in US 5,252,389.

With the incorporation of clean-polypropylene into component (B) of the dielectric layer in particular the attenuation behaviour of said layer is still further improved.

It is preferred that component (B) of the dielectric layer comprises at least 50 wt% of clean-polypropylene.

In a further preferred embodiment, the ratio of components (A):(B) of the dielectric layer of the inventive coaxial cable is from 1:99 to 60:40, more preferably from 25:75 to 60:40.

Further preferred, the dielectric layer of the inventive coaxial cable has been expanded.

Expansion can be performed via chemical foaming in which the polymer raw material used for the dielectric layer is compounded with a chemical foaming agent which on decomposition blows closed cells of desired size into the dielectric layer. However, preferably expansion is achieved by physical foaming in which during extrusion of the dielectric material inert gas such as nitrogen, carbon dioxide or argon is injected to blow gas filled expanded cells.

It is preferred that the degree of expansion in the dielectric layer is at least 60 vol%, more preferred at least 75 vol% and most preferred between 77 and 85 vol%.

Furthermore, it is preferred that the dielectric layer of the inventive coaxial cable further comprises a nucleating agent, preferably in an amount of 0.01 to 0.05 wt%.

As the improved properties of the inventive coaxial cable in particular show up at higher radio frequencies it is preferred that the coaxial cable is used for the transmission of electromagnetic signals with a frequency of above 1 GHz, more preferably of above 1.5 GHz.

As mentioned, the present invention also relates to the use of propylene homo- or copolymer having strain hardening behaviour for the production of a dielectric layer of a coaxial cable.

In the following the present invention will further be illustrated by means of examples with reference to the Figures:

- Fig. 1: shows a typical coaxial cable design comprising an inner conductor (1), an inner skin/adhesion layer (2), a foamed dielectric (3), an outer skin (4), an outer conductor (5) and a jacket (6);
- Fig. 2: shows a schematic drawing of the apparatus used for determining strain hardening behaviour as well as a schematic diagram resulting from the measurement, and
- Fig. 3: shows a diagram showing recorded melt strength vs. drawability curves of different polymers with and without strain hardening behaviour.
- Fig. 4: shows a diagram showing melt strength vs. drawability curves of several polymers and polymer blends as used in the following examples.

Examples

1) Definition and measurement of strain hardening behaviour

The term "strain hardening behaviour" as used herein is defined according to Fig. 2 and 3. Fig. 2 shows a schematic representation of the experimental procedure which is used to determine strain hardening.

The strain hardening behaviour of polymers is analysed by Rheotens apparatus 7 (product of Göttfert, Siemensstr. 2, 74711 Buchen, Germany) in which a melt strand 8 is elongated by drawing down with a defined acceleration. The haul-off force F in dependence of draw-down velocity v is recorded.

The test procedure is performed in a standard acclimatised room with controlled room temperature of T = 23°C. The Rheotens apparatus 7 is combined with an extruder/melt pump 9 for continuous feeding of the melt strand 8. The extrusion temperature is 200°C; a capillary die with a diameter of 2 mm and a length of 6 mm is used and the acceleration of the melt strand 8 drawn down is 120 mm/s^2 .

The schematic diagram in Fig. 2 shows in an exemplary fashion the measured increase in haul-off force F (i.e. "melt strength") vs. the increase in draw-down velocity v (i.e. "drawability").

Figure 3 shows the recorded curves of Rheotens measurements of polymer samples with and without strain hardening behaviour. The maximum points $(F_{max}; v_{max})$ at failure of the strand are characteristic for the strength and the drawability of the melt. The standard unmodified propylene polymers 10, 11 and 12 with melt flow rates of 0.3, 2.0 and 3.0 g/10 min at 230°C/2.16 kg, respectively, show a very low melt strength and low drawability. Accordingly, they have no strain hardening behaviour.

Modified propylene polymers 13 (melt flow rate of sample in diagram is 2 to 3 g/10 min at 230°C/2.16 kg) or LDPE 14 (melt flow rate of sample in dia-

gram is 0.7 g/10 min at 230°C/2.16 kg) show a completely different melt strength vs. drawability behaviour:

With increasing the draw down velocity v the haul-off force F increases to a much higher level, compared to the standard propylene polymers 10, 11 and 12. This curve shape is characteristic for strain hardening behaviour. While polymers 10 and 11 show haul-off F_{max} larger than 5cN, they do not have strain hardening behaviour because they do not have draw-down velocities v_{max} larger than 150 mm/s.

Accordingly, propylene polymers which have strain hardening behaviour as used herein have enhanced strength with haul-off forces $F_{max} > 5$ cN and enhanced drawability with draw-down velocities $v_{max} > 150$ mm/s.

2) Synthesis of propylene homopolymer with strain hardening behaviour

A powdery polypropylene homopolymer, with a melt index of 0.25 g/10 min at 230°C/2.16 kg and an average particle size of 0.45 mm, is metered continuously into a continuous mixer. Furthermore, 0.45 wt% based on the propylene homopolymer of tert.-butyl peroxybenzoate as thermally decomposing free radical forming agent is metered into the mixer. While being mixed homogeneously at 50°C, the propylene homopolymer containing the tert.butyl peroxybenzoate is charged absorptively during a residence time of 7 minutes at 50°C by means of a mixture of butadiene and nitrogen with 0.135 wt% of butadiene, based on the polypropylene homopolymer. After transfer to a twin screw extruder, the powdery reaction mixture, in contact with the mixture of butadiene and nitrogen, with which it has been charged, is melted at a mass temperature of 230°C and, after a coarse degassing, subjected to a fine degassing with addition of water as an entraining agent, an additive mixture of 0.1 wt% of tetrakis-(methylene-(3,5-di-t-butylhydroxycinnamate)-methane, 0.1 wt% of tris-(2,4-di-t-butylphenyl)-phosphite), 0.1 wt% of pentaerythritol tetrakis-3-(3,5-di-t-butyl-4-hydroxyphenyl)-propionate and 0.1 wt% of calcium stearate is added to the melt. After distribution of additives the melt is discharged and granulated.

The resulting, modified propylene polymer MPP shows strain hardening behaviour characterized by the Rheotens values of $F_{max} = 38$ cN and $v_{max} = 175$ mm/s measured at failure of the strand and a melt index of 2.3g/10 min at 230° C/2.16 kg.

From figure 4 it can be seen that MPP shows similar strain hardening behaviour as LDPE, and MDPE/HDPE show similar behaviour as clean PP.

3) Measurement of electronic properties

For measuring the electronic properties, square specimens with 9 cm x 9 cm dimensions and a thickness of 4.0 mm were produced by press moulding of the polymer compositions with 15°C/min cooling in accordance to ISO 293 - 1986 (E).

The dielectric properties (dissipation, relative permittivity) have been measured using the split post resonator technique at a nominal frequency of 1.8 GHz.

Density as given in Table 1 was measured according to ISO 1872-2-B/ISO 1183D. Melt flow rate was measured according to ISO 1133 at a load of 2.16 kg at 230°C for all polymer materials (PP and PE).

From Table 1 it can be seen that a mixture of MDPE + 25 wt% LDPE has a dissipation factor of 118 whereas a blend of clean-PP and 25 wt% MPP shows a strongly reduced dissipation factor of 77.

Table 1: Electrical measurements at high frequency

Polymer composition	Density (kg/m³)	MFR ₂ ^{230°C}	Dissipation factor Tan Delta at 1.8 GHz	Relative permit- tivity Epsilon at 1800 GHz
LDPE	923	6	163	2,29
MDPE	936	4.8	116	2,32
HDPE	952	5.3	102	2,35
MDPE + 25% LDPE	932	5	118	2,3
HDPE + 25% LDPE	946	5.5	96	2,33
·	Examples acco	ording to the ir	nvention	
MPP	910	2.5	128	2,26
Clean PP	910	3.7	60	2.25
15 wt% MPP + clean PP	910	3.5	69	2.24
25 wt% MPP + clean PP	910	3.4	77	2.25
35 wt% MPP + clean PP	910	3.3	86	2.23
45 wt% MPP + clean PP	910	3.2	95	2.25

Claims

- 1. A coaxial or triaxial cable comprising a dielectric layer which comprises as a component (A) a propylene homo- or copolymer having strain hardening behaviour.
- 2. Cable according to claim 1, wherein the dielectric layer further comprises as a component (B) a medium or high density ethylene homo- or copolymer and/or a non-strain hardening behaviour propylene homo- or copolymer.
- 3. Cable according to claim 2, wherein component (B) comprises a propylene homo- or copolymer having a catalyst residue of less than 50 ppm, an ash content below 100 ppm and a chloride content of less than 5 ppm.
- 4. Cable according to claim 3, wherein the propylene homo-or copolymer is having a catalyst residue of less than 5 ppm, an ash content below 30 ppm, and a chloride content of less than 1 ppm.
- 5. Cable according to any of claims 3 and 4 wherein component (B) comprises at least 50 wt% of said polypropylene.
- 6. Cable according to any of the preceding claims, wherein the ratio of components (A):(B) is from 1:99 to 60:40, more preferably from 25:75 to 60:40.
- 7. Cable according to any of the preceding claims wherein the propylene homo- or copolymer having strain hardening behaviour has a melt flow rate of 0.1 to 25 g/10min at 230°C/2.16kg.
- 8. Cable according to any of the preceding claims wherein the dielectric layer has been expanded, preferably by physical foaming.
- 9. Cable according to claim 8, wherein the degree of expansion is at least 60%, more preferably at least 75%.

- 10. Cable according to any of the preceding claims wherein the dielectric layer further comprises a nucleating agent, preferably in an amount of 0.01 to 0.05 wt%.
- 11. Use of propylene homo- or copolymer having strain hardening behaviour for the production of a dielectric layer of a coaxial or triaxial cable.

Fig. 1

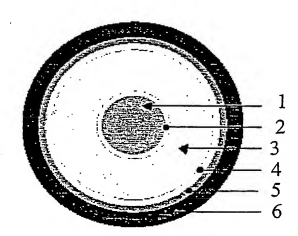


Fig. 2

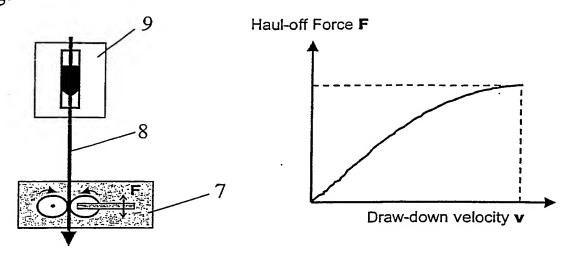


Fig. 3

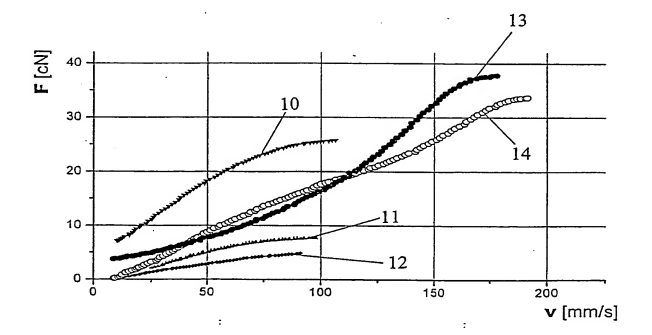
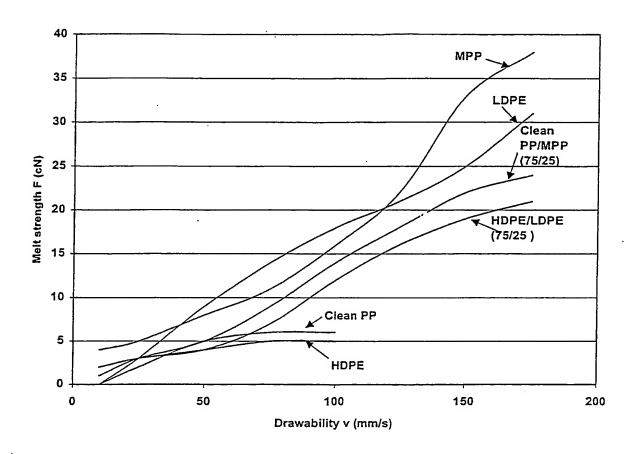


Fig. 4



INTERNATIONAL SEARCH REPORT

			PCT/E 03	3/11905
	IFICATION OF SUBJECT MATTER H01B3/44			
1166 /	HU163/44			
According to	o International Patent Classification (IPC) or to both national classific	cation and IRC		
B. FIELDS	SEARCHED			
Minimum de IPC 7	ocumentation searched (classification system followed by classifical $H01B$	lion symbols)		
/				
Documenta	tion searched other than minimum documentation to the extent that	such documents are inclu	ided in the fields s	earched
Electronic d	lata base consulted during the international search (name of data ba	ase and, where practical,	search terms used	i)
EPO-In	ternal, WPI Data, PAJ			
i F				
0.000114	ENTS CONSIDERED TO DE DEL SIA AR			
Category °	ENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the re	levani passages		Relevant to claim No.
Х	EP 0 961 295 A (UNION CARBIDE CH			1-11
	PLASTIC) 1 December 1999 (1999-12 page 5, paragraph 22; claim 1	2-01)		
χ	US 3 968 463 A (BOYSEN ROBERT L)			1-11
^	6 July 1976 (1976-07-06)			1-11
	claims 1-5			
Х	DE 14 65 640 B (KABEL METALLWERKE	E GHH)		- 1-11
	6 November 1969 (1969-11-06) claims 1,2			
v		EDICON CDA		
Х	CH 357 096 A (ZIEGLER KARL ;MONTE (IT)) 30 September 1961 (1961-09-			1-11
	claim I			
	-	-/		
				•
X Funt	ner documents are listed in the continuation of box C.	χ Patent family n	nembers are listed	in annex.
° Special cal	legories of cited documents:	*T' later document publi		
consid	nt defining the general state of the art which is not ered to be of particular relevance	or priority date and cited to understand invention		
filing d	ale	'X' document of particul cannot be consider	ed novel or cannot	be considered to
which i	nt which may throw doubts on priority claim(s) or is cited to establish the publication date of another I or other special reason (as specified)	"Y" document of particul	ar relevance; the c	cument is taken alone laimed invention ventive step when the
other n		document is combit ments, such combit	ned with one or mo	re other such docu- is to a person skilled
	nt published prior to the international filling date but an the priority date claimed	in the art. "&" document member of	of the same patent t	amily
Date of the a	actual completion of the international search	Date of mailing of the	ne international sea	rch report
22	2 January 2004	29/01/20	004	
Name and m	nailing address of the ISA E: _paan Patent Office, P.B, 5818 Patentlaan 2	Authorized officer		
	NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl.	Lehnert,	Δ	
	Fax: (+31-70) 340-3016	Leimei C,		

Form PCT/ISA/210 (second sheet) (July 1992)



international Application No PCT/L 03/11905

on) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
US 6 121 335 A (HIGASHIKUBO TAKASHI ET AL) 19 September 2000 (2000-09-19) cited in the application column 7, line 8 - line 12 column 7, line 23 - line 30	1-11
US 6 130 385 A (TUUNANEN VESA ET AL) 10 October 2000 (2000-10-10) cited in the application column 6, line 11 - line 12	1-11
EP 0 885 918 A (TORAY INDUSTRIES) 23 December 1998 (1998-12-23) example 1	1-11
EP 0 190 889 A (HIMONT INC) 13 August 1986 (1986-08-13) cited in the application page 12, line 27 - line 32	1-11
EP 0 384 431 A (HIMONT INC) 29 August 1990 (1990-08-29) cited in the application the whole document	1-11
	column 7, line 8 - line 12 column 7, line 23 - line 30 US 6 130 385 A (TUUNANEN VESA ET AL) 10 October 2000 (2000-10-10) cited in the application column 6, line 11 - line 12 EP 0 885 918 A (TORAY INDUSTRIES) 23 December 1998 (1998-12-23) example 1 EP 0 190 889 A (HIMONT INC) 13 August 1986 (1986-08-13) cited in the application page 12, line 27 - line 32 EP 0 384 431 A (HIMONT INC) 29 August 1990 (1990-08-29) cited in the application

INTERNATIONAL SEARCH REPORT

In atlon on patent family members

PCT/E 03/11905

	atent document d in search report		Publication date		Patent family member(s)		Publication date
EP	0961295	A	01-12-1999	US BR CA EP	6599626 9901394 2272737 0961295	A A1	29-07-2003 18-09-2001 26-11-1999 01-12-1999
US	3968463	А	06-07-1976	AU AU CA DE FR GB IT JP NL SE BE	499769 7206774 1020644 2437998 2240507 1477874 1019782 50072185 7410616 7410134 818568	A A1 A1 A B A A	03-05-1979 12-02-1976 08-11-1977 06-03-1975 07-03-1975 29-06-1977 30-11-1977 14-06-1975 11-02-1975 07-03-1975
DE	1465640	В	06-11-1969	NONE			
СН	357096	Α	30-09-1961	NONE			
US	6121335	A	19-09-2000	JP JP BE CN DE	3457543 2000072912 1013525 1246492 19940604	A A3 A ,B	20-10-2003 07-03-2000 05-03-2002 08-03-2000 02-03-2000
US	6130385	A	10-10-2000	FI AT AU BR CA CN DE DE DK EP ES WO JP KR NO PT	962715 225560 3346497 9710189 2258317 1224528 69716073 69716073 909449 0909449 2184104 9801870 2000512796 2000022474 986167 909449	T A A A1 A ,B D1 T2 T3 A1 T3 A1 T A	02-01-1998 15-10-2002 02-02-1998 10-08-1999 15-01-1998 28-07-1999 07-11-2002 13-03-2003 30-12-2002 21-04-1999 01-04-2003 15-01-1998 26-09-2000 25-04-2000 28-12-1998 28-02-2003
EP	0885918		23-12-1998	JP JP CA DE DE EP US CN WO ID	10156938 10112419 2229778 69714022 69714022 0885918 6094337 1198759 9806776 19278 5275698	A A1 D1 T2 A1 A A,B A1 A	16-06-1998 28-04-1998 19-02-1998 22-08-2002 28-11-2002 23-12-1998 25-07-2000 11-11-1998 19-02-1998 28-06-1998 12-11-1998
EP	0190889	Α	13-08-1986	AT AU	88196 616106		15-04-1993 17-10-1991

INTERNATIONAL SEARCH REPORT

Infamotion on patent family members

International Application No PCT/E. 03/11905

Patent document cited in search report	Publication date		Patent family member(s)	Publication date
EP 0190889 A		ΑU	5215890 A	19-07-1990
		AU	597241 B2	31-05-1990
		AU	5284386 A	07-08-1986
		ΑU	7942891 A	12-09-1991
		BR	8600413 A	14-10-1986
		CA	1280543 C	19-02-1991
		CN	86100791 A ,B	04-02-1987
		CN	1041769 A ,B	02-05-1990
		DE	3688258 D1	19-05-1993
		DE	3688258 T2	29-07-1993
		EP	0190889 A2	13-08-1986
		IN	166935 A1	11-08-1990
		IN	170549 A1	11-04-1992
		JP	7045551 B	17-05-1995
		JP	62121704 A	03-06-1987
		NO	860330 A	01-08-1986
		NO	167039 B	17-06-1991
		PT	81945 A ,B	01-02-1986
		RU	2031906 C1	27-03-1995
		US	5731362 A	24-03-1998
		US	4916198 A	10-04-1990
		ÜS	5554668 A	10-09-1996
		US	5591785 A	07-01-1997
		ZA	8600528 A	24-09-1986
EP 0384431 A	29-08-1990	US	5047485 A	10-09-1991
		ΑT	114677 T	15-12-1994
		AU	616103 B2	17-10-1991
		ΑU	4994390 A	30-08-1990
		BR	9000791 A	22-01-1991
		CA	1339476 C	23-09-1997
		CN	1045107 A ,B	05-09-1990
		DE	69014348 D1	12-01-1995
		DE	69014348 T2	20-04-1995
		DK	384431 T3	16-01-1995
	•	EP	0384431 A2	29-08-1990
		ES	2064503 T3	01-02-1995
		JP	2298536 A	10-12-1990
		JP	2744317 B2	28-04-1998
		KR	145323 B1	15-07-1998
		NO	900738 A	22-08-1990
		PT	93206 A	31-08-1990
		RU	2036931 C1	09-06-1995
		ZA	9000911 A	28-12-1990